Appl. No. 09/783,253 Amdt. dated June 27, 2003 Amendment under 37 CFR 1.116 Expedited Procedure Examining Group

This listing of claims will replace all prior versions, and listings, of claims in the application:

## **Listing of Claims:**

Please amend Claims 78, 102-106, and 110, to read as follows. Please cancel Claims 1-77 and Claims 80-101.

Claims 1-77 (cancel).

78. (currently amended) A method for laminal substance delivery, said method comprising:

providing a luminal prosthesis incorporating and/or coupled to the substance, wherein the prosthesis eontains comprises a rate limiting barrier; and implanting the prosthesis in a body lumen so that the substance is released from the prosthesis at multiple rates including an initial rate and at least one subsequent rate which is substantially higher than the initial rate and substantial substance release from the barrier begins after a an appreciable preselected time period.

79. (original) A method as in claim 78, wherein the barrier has a sufficient thickness to allow diffusion of the substance through the barrier.

Claims 80-101 (cancel).

- 102. (currently amended) A method as in any of claims 78-79, wherein substantial release of the substance the at least one subsequent rate begins within a time period of ranging from 4 hours to 24 weeks in a vascular environment.
- 103. (currently amended) A method as in any of claims 102, wherein substantial release of the substance the at least one subsequent rate begins within a time period of 1 day to 12 weeks in a vascular environment.
- 104. (currently amended) A method as in any of claims 102, wherein substantial release of the substance the at least one subsequent rate begins within a time period of 2 days to 8 weeks in a vascular environment.

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- 105. (currently amended) A method as in any of claims 102, wherein substantial release of the substance the at least one subsequent rate begins within a time period of 3 days to 50 days in a vascular environment.
- 106. (currently amended) A method as in any of claims 78—79 or 102—105:, further comprising directing energy at the prosthesis to effect release of the substance from the prosthesis.
- 107. (previously added) A method as in claim 78, wherein the prosthesis incorporates the substance by coating, spraying, dipping, deposition or painting the substance on the prosthesis.
- 108. (previously added) A method as in claim 18, wherein the substance is incorporated in a reservoir in or on a scaffold containing the substance.
- 109. (previously added) A method as in claim 106, wherein the energy is at least one of ultrasound, magnetic resonance imaging, magnetic field, radio frequency, temperature change, electromagnetic, x-ray, radiation, hear, gamma, or microwave.
- 110. (currently amended) A method as in claim 78–79 or 102–105 wherein the prosthesis incorporates magnetic particles coupled to the substance and further comprising the step of directing a magnetic field at the prosthesis to effect release of the substance from the prosthesis.
- 111. (previously added) A method as in any one of Claims 78-79 wherein the substance comprises at least one agent selected from the consisting of immunosuppressant agent, anti-inflammatory agent, anti-proliferative agent, anti-migratory agent, anti-fibrotic agent, anti-thrombotic agent, anti-platelet agent, and IIb/IIIa agent.